AMENDMENTS TO THE SPECIFICATION:

Please delete the previously submitted Sequence Listing and insert the attached Sequence Listing (pages 1-19) at the end of the specification.

Please amend the Brief Description of Drawings on page 4, line 23-24, as follows:

BRIEF DESCRIPTION OF DRAWINGS

Fig. 1 shows an alignment of lipase sequences, as follows.

"rhimi" shows the Rhizomucor miehei lipase (SEQ ID NO:30)

"rhid!" shows the Rhizomucor delemar lipase (SEQ ID NO:31)

"SP400" shows the Thermomyces (Humicola) lanuginosus lipase (SEQ ID NO:32)

"Pcl" shows the Penicillium camemberti lipase (SEQ ID NO:33)

"FoLnp11" shows the Fusarium oxysporum lipase (SEQ ID NO:34)

Please amend the paragraph on page 7, lines 1-6, as follows:

For the Rhizomucor miehei lipase, the extent of the alcohol binding site can be found from the PDB file "5tgl.pdb" available in Structural Classification of Proteins (SCOP) on the Internet, at https://www.rcsb.org/pdb/, showing the complex with the inhibitor n-hexylphosphonate ethyl ester which mimics the substrate. It is described in Derewenda et al. (supra), Brzozowski et al. (supra) and Brady et al. (supra). The sn2 position of this model is the atom CE2.

Please amend the paragraph on page 7, lines 28-33, as follows:

The procedure is conveniently performed by use of software such as MSI's Insight II. It involves alignment with 4TGL, a crystal structure of the lipase from *Rhizomucor miehei* inhibited irreversibly by diethyl p-nitrophenyl phosphate. This is available in Structural Classification of Proteins (SCOP) on the Internet, at http://www-rcsb.org/pdb/, and is described in Derewenda et al. (supra). The *Rhizomucor miehei* lipase comprises a catalytic triad consisting of the amino acid residues \$144, D203 and H 257.

Please amend the paragraph on page 8, lines 22-27, as follows:

Alternatively, the variant may carry a peptide extension at the C-terminal and/or the N-terminal. The C-terminal extension may consist of 1-10 amino acid residues, e.g. A, P, AG, DG, PG, AGG, PVGF (SEQ ID. NO. 15), AGRF (SEQ ID. NO. 16), PRGF (SEQ ID. NO. 17). AGGF (SEQ ID. NO. 18) or AGGFS (SEQ ID. NO. 19); or it may consist of 40-50 residues, e.g., consisting of the 48 C-terminal residues of the Fusanium oxysporum lipase GGFSWRRYRSAESVDKRATMTDAELEKKLNSVVQMD KEYVKNNQARS (SEQ ID. NO. 20). The C-terminal extension may increase the phospholipase activity.

Please amend the paragraph on page 10, lines 3-7, as follows:

The variant of the invention may further comprise a peptide extension at the N-terminal, e.g. consisting of 1-15 (particularly 4-10) amino acid residues, and specifically comprising 1, 2 or 3 positively charged amino acids. Some specific N-terminal peptide extensions are AS, SPIRR (SEQ ID. NO. 21). E1RP, E1SPIRPRP (SEQ ID. NO. 22), E1SPPRRP (SEQ ID. NO. 23) and E1SPIRPRP (SEQ ID. NO.22), Further, any peptide extension described in WO 97/04079 and WO 97/07202 may be used.

Please amend the paragraph on page 11, lines 1-7, as follows:

Specific variants of the *H. lanuginosa* lipase are disclosed in the examples. Corresponding alterations may be made in other parent lipolytic enzymes. Further variants may be derived from these by omitting amino acid modifications at positions 1, 106, 186, 225, 232, 237, 239 or 274. Variants with 274S may optionally have a further C-terminal extension of WRRYRSAESVDKRATMTDAELEKKLNSYVOM DKEYVKNNQARS (SEQ. ID. NO. 24) (corresponding to the C-terminal of the *F. oxysporum* lipase) in full or truncated form.

Please amend the paragraph on page 11, lines 13-18, as follows:

T267stop indicates a stop codon, i.e. deletion of T267 and all following amino acids (i.e. C268 and L269). 270P, 271V indicates a C-terminal extension of PV (i.e. at new positions 270 and 271). -G268 indicates deletion of G at position 266. Parentheses indicate that the alteration is optional, or in examples that the alteration is uncertain. SPIRR (SEQ ID NO:21) indicates an

N-terminal extension. D266 may refer to the position or to substitution with any amino acid (except D).

Please amend the paragraph on page 11, lines 19-22, as follows:

E1SPPCGRRP (SEQ ID. NO. 25) or SPPCGRRP(-E) (SEQ ID NO.25) indicates a substitution of E1 with SPPCGRRP (SEQ ID NO.25), i.e. a peptide addition at the N-terminal. T267GS indicates a substitution of T267 with GS, or in other words the substitution T267G and an insertion of S between G267 and C268.

Please amend the paragraph on page 17, lines 17-20, as follows:

Specific commercially available protease enzymes include Alealase . Savinase . Primase . Duralase . Esperase . and Kannase . (Nove Nordisk A/S), Maxatase . Maxacal . Maxapam . Properses . Purafect . Purafect . OxP . FN2 . And . FN3 . And . Commercial Inc.) ALCALASE. SAVINASE. PRIMASE. DURALASE. SPERASE . and KANNASE . (Novozymes A/S), MAXATASE. MAXACAL . MAXAPEM . PROPERASE . PURAFECT. PURAFECT . OxP . FN2 . And FN3 . (Genencor International Inc.).

Please amend the paragraph on page 17, lines 32-35, as follows:

Commercially available cellulases include Celluzyme*, and Carezyme* (Novo Nordisk A/S), Clazinase*, and Puradax HA* (Genencer International Inc.), and KAC-500(B)* (Kac Corporation)CELLUZYME and CAREZYME* (Novozymes A/S), CLAZINASE*, and PURADEX HA* (Genencer International Inc.), and KAC-500(B)* (Kac Corporation)

Please amend the paragraph on page 27, lines 14-18, as follows:

In this SLU method, the lipase activity is measured at 30—°C and pH 9 with a stabilized olive oil emulsion (Sigma catalog No. 800-1) as the substrate, in a 5 mM Tris buffer containing 40 mM NaCl and 5 mM calcium chloride. 2.5 ml of the substrate is mixed with 12.5 ml buffer, the pH is adjusted to 9, 0.5 ml of diluted lipase sample is added, and the amount of oleic acid formed is followed by titration with a pH stat.

Please amend the paragraph on page 27, lines 19-20, as follows:

One SLU is the amount of lipase which liberates 1 —µmole of titratable oleic acid per minute under these conditions.

Please amend the paragraph on page 31, lines 7-18, as follows:

The gene encoding the lipolytic enzyme in question is inserted into the plasmid pHD414. In accordance with the manufacturer's instructions the Scal site of the Ampicillin gene of pHD414 is changed to a Mlul site by use of the following primer:

Primer 3: AGAAATCGGGTATCCTTTCAG. (SEQ I.D. NO. 27)

The pHD414 vector comprising the lipolytic gene in question is then used as a template for DNA polymerase and oligos 7258 and 7770.

7258: 5'p gaa tga ctt ggt tga cgc gtc acc agt cac 3' (SEQ I.D. NO. 28)

(Thus changing the Scal site found in the ampicillin resistance gene and used for cutting to a Miul site).

Primer no. 7770 was used as the selection primer.

7770: 5'p tot ago coa gaa tac tgg atc aaa to 3'(<u>SEQ I.D. NO 29</u>) (Changes the Scal site found in the *H. lanuginosa* lipase gene without changing the amino acid sequence).

Please amend the paragraph on page 33, lines 32-35 as follows:

SPIRR (<u>SEQ ID NO:21</u>), G91A, D96W, E99K, W260C, G263Q, L264A, I265T, G266D. T267A, L269N, 270A, 271G, 272, G273F, (274S)

SPIRR (<u>SEQ ID NO:21)</u>, G91A, D96W, E99K, G263Q, L264A, I265T, G266D, T267A, L269N, 270A, 271G, 272G, 273F, (274S)

Please amend the paragraph on page 35, lines 1-4, as follows:

SPIRR (<u>SEQ ID NO:21)</u>, D96W, E99K, G263Q, L264A, I265T, G266D, T267A, L269N, 270A, 271G, 272G, 273F, (274S)

SPIRR (<u>SEQ ID NO:21)</u>, G91A, D96W, G263Q, L264A, I265T, G266D, T267A, L269N, 270A, 271G, 272G, 273F, (274S)

Please amend the table under Example 5, beginning on page 37, as follows:

E1A, G91A, D96W, E99K, P256A, W260H, G263Q, L264A, I265T, G266D, T267A, L269N 270A, 271G, 272G, 273F, (274S)
SPIRR (<u>SEQ ID NO:21</u>), G91A, D96W, E99K, G263Q, L264A, I265T, G266D, T267A, L269N 270A, 271G, 272G, 273F, (274S)
E1A, G91A, D96W, P256A, W260H, G263Q, L264A, I265T, G266D, T267A, L269N, 270A 271G, 272G, 273F, (274\$)
E1A, G91A, D96W, E99K, P256A, W260H, G263Q, L264A, I265T, G266D, T267A, L269N
E1A, G91A, D96W, E99K, Q249R, G266S, 270D, 271G
E1A, G91A, D96W, E99K, Q249R, G266D
E1A, G91A, D96W, E99K, Q249R, G266A, 270P, 271G
G266D
E1SPPCGRRP (SEQ ID NO:25) +E99N +E239C +Q249R +G266D
E1SPPCGRRP (SEQ ID NO:25) +E239C +Q249R +G266D
E1SPPCGRRP (SEQ ID NO:25) +L93K +E99K +E239C +Q249R +G266D
E1SPPCGRRP (SEQ ID NO:25) +E99K +E239C +Q249R +G266D
G266A
G266W
G266V
G263Q +L264A +I265T +G266D +T267A
G263F +L264A +G266S +T267E
E1SPPCGRRP (SEQ ID NO:25) +E239C +Q249R +G263Q +L264A +I265T +G266D +T267A
G266S
G266L
G263A +G266A
G263A +G266Y
E1SPPCGRRP (SEQ ID NO:25) +E239C +Q249R +G266A
E1SPPCGRRP (SEQ ID NO:25) +E239C +Q249R +G266S
E1SPPCGRRP (SEQ ID NO:25) +E239C +Q249R +G263F +L264A +G266S +T267E
D62A + G266A
D62A + G266S
D96S + G266A
D96S+ G266S

D96S+ G266R

D96S+ G266W

D96S+ G266V

E1SPPCGRRP (SEQ ID NO:25) + G91A+ D96W+ E239C+ Q249R+ G266D

E1SPPCGRRP (SEQ ID NO:25) + G91A+ D96W+ E239C+ Q249R+ G266S

E1SPPCGRRP (SEQ ID NO:25) + G91A+ D96W+ E239C+ Q249R+ G263E+ G266S+ 270A

E1SPPCGRRP (SEQ ID NO:25) + G91A+ D96W+ E239C+ Q249R+ L264P+ G266S

E1SPPCGRRP (SEQ ID NO:25) + G91A+ D96W+ E239C+ Q249R+ P256T+ G266D

E1SPPCGRRP (SEQ ID NO:25) + G91A+ D96W+ E239C+ Q249R+ G266C+ T267P+ L269stop

G263D +L264I +I265N +G266E +T267GS

E219G +L264I +I265N +G266T +T267GL

E1A+ G91A+ D96W+ E99K+ P256A+ W260H+ G263Q+ L264A+ I265T+ G266D+ T267A+ L269N+ 270A+ 271G+ 272G+ 273F (+274S)

E1A+ G91A+ D96W+ E99K+ E239C+ Q249R+ P256A+ G263Q+ L264A+ I265T+ G266D+ T267A+ L269N+ 270A+ 271G+ 272G +273F (+274S)

E1A+ G91A+ D96W+ E99K+ N248T+ Q249R+ W260Q+ G263Q+ L264A+ I265T+ G266D+ T267A+ L269N+ 270A+ 271G+ 272G+ 273F (+274S)

SPIRR (<u>SEQ ID NO:21)</u> + G91A+ D96W+ E99K+ W260C+ G263Q+ L264A+ I265T+ G266D+ T267A+ L269N+ 270A+ 271G+ 272+ G273F (+274S)

SPIRR (<u>SEQ ID NO:21</u>) + G91A+ D96W+ E99K+ G263Q+ L264A+ I265T+ G266D+ T267A+ L269N+ 270A+ 271G+ 272G+ 273F (+274S)

E1A+ G91A+ D96W+ E99K+ G263Q+ L264A+ I265T+ G266D+ T267A+ L269N+ 270A+ 271G+ 272G+ 273F (+274S)

E1A+ G91A+ D96W+ E99K+ P256A+ W260H+ G263Q+ L264A+ I265T+ G266D+ T267A+ L269N+ 270A+ 271G +272G +273F (+274S)

SPIRR (<u>SEQ ID NO:21</u>) + D96W+ E99K+ G263Q+ L264A+ I265T+ G266D+ T267A+ L269N+ 270A+ 271G+ 272G+ 273F (+274S)

SPIRR (<u>SEQ ID NO:21</u>) + G91A+ D96W+ G263Q+ L264A+ I265T+ G266D+ T267A+ L269N+ 270A+ 271G+ 272G+ 273F (+274S)

E1A+ G91A+ D96W+ E99K+ P256A+ W260H+ G263Q+ L264A+ I265T+ G266D+ T267A+ L269N

E1A+ G91A+ D96W+ E99K+ Q249R+ G263E+ G266D+ L269N+ 270P+ 271V+ 272G+ 273F

E1A+ G91A+ D96W+ E99K+ Q249R+ G263A+ G266S+ L269N+ 270A+ 271G+ 272R+ 273F

E1A+ G91A+ D96W+ E99K+ Q249R+ L264P+ A266+ L269I+ 270P+ 271R+ 272G+ 273F

E1A+ G91A+ D96W+ E99K+ Q249R+ L264C+ I265N+ G266P+ T267stop

E1A+ G91A+ D96W+ E99K (+R232L)+ Q249R+ G266S+ 270A

E1A+ G91A+ D96W+ E99K+ Q249R+ G266S+ 270D+ 271G F1A+ G91A+ D96W+ F99K+ Q249R+ L264F+ A266+ 270A+ 271G+ 272G+ 273F E1A+ G91A+ D96W+ E99K+ Q249R+ L264G+ I265G+ G266F+ T267stop E1A+ G91A+ D96W+ E99K+ Q249R+ L264stop E1A+ G91A+ D96W+ E99K+ P256A+ W260H+ G263Q+ L264A+ I265T+ G266D+ T267A+ L269N+ 270A+ 271G E1A+ G91A+ D96W+ E99K+ P256A+ W260H+ G263Q+ L264A+ I265T+ G266D+ T267A+ L269N+ 270A+ 271G+ 272G E1A+ G91A+ D96W+ E99K+ Q249R+ G266D E1A+ G91A+ D96W+ E99K+ Q249R+ G266D E1A+ G91A+ D96W+ E99K+ Q249R+ G266A+ 270P+ 271G E1A+ G91A+ D96W+ E99K+ Q249R+ L264P+ I265F+ L269stop E1A+ G91A+ D96W+ E99K+ Q249R+ G266D+ L269S+ 270A+ 271G+ 272G+ 273F E1A+ G91A+ D96W+ E99K+ Q249R+ G266D+ L269N+ 270A E1A+ G91A+ D96W+ E99K+ Q249R+ G266S+ L269N+ 270A+ 271G+ 272G+ 273F E1A+ G91A+ D96W+ E99K+ Q249R+ L264P+ L267Q+ L269N E1A+ G91A+ D96W+ E99K+ Q249R+ G263R+ I265L+ L269N+ 270P E1A+ D96W+ E99K+ P256A+ W260H+ G263Q+ L264A+ I265T+ G266D+ T267A+ L269N+ 270A+ 271G+ 272G+ 273F (+274S) E1A+ G225R+ G266D E1A+ G225R+ G263A+ I265V+ G266S F1A+ G225R+ G263A+ T267A E1SPPCGRRP (SEQ ID NO:25) + D96S+ E239C+ Q249R+ I252M+ L264Q+ G266D E1SPPCGRRP (SEQ ID NO:25) + G91A+ D96W+ E239C+ Q249R+ G266D E1SPPCGRRP (SEQ ID NO:25) + D96S+ E239C+ Q249R+ G266D E1SPPCGRRP (SEQ ID NO:25) + D96S+ E239C+ Q249R+ G266C+ L267A E1A+ G91A+ D96W+ E99K+ Q249R+ G266A E1A+ D96M+ G106S+ G225R+ G266D E1A+ D96Q+ G106S+ G225R+ G266S E1A+ D96F+ G225R+ G266S E1A+ D96C+ G225R+ G266T

E1A+ D96H+ G106S+ G225R+ G266S SPIRR (<u>SEQ ID NO:21</u>) + D96S+ G266D SPIRR (<u>SEQ ID NO:21</u>) + D96R+ G106S+ G266D SPIRR (SEQ ID NO:21) + D96I+ G106S+ G266S SPIRR (SEQ ID NO:21) + D96W+ K237R+ G266\$ SPIRR (SEQ ID NO:21) + G266A SPIRR (SEQ ID NO:21) + D96S+ G106S+ G225R+ G266D SPIRR (SEQ ID NO:21) + D96Q+ G106S+ G225R+ G266A SPIRR (SEQ ID NO:21) + D96Y+ G106S+ G225R+ G266N SPIRR (SEQ ID NO:21) + D96C+ G106S+ G225R+ G266T SPIRR (SEQ ID NO.21) + D96H+ T186I+ G225R+ G266S E1SPPRRP (SEQ ID NO:23) + G91A+ D96W+ E239C+ O249R+ G266D E1SPPRRP (SEQ ID NO:23) + G91A+ D96W+ E239C+ Q249R+ G266S E1SPPRRP (SEQ ID NO:23) + G91A+ D96W+ E239C+ Q249R+ G263E+ G266S+ 270A E1SPPRRP (SEQ ID NO:23) + G91A+ D96W+ E239C+ Q249R+ L264P+ G266S E1SPPRRP (SEQ ID NO:23) + G91A+ D96W+ E239C+ Q249R+ P256T+ G266D E1SPPRRP (SEQ ID NO:23) + G91A+ D96W+ E239C+ Q249R+ G266C+ T267P+ L269stop E1A+ G91A+ D96W+ E99K+ Q249R+ G266S+ T267S E1SPPCGRRP (SEQ ID NO:25) + G91A+ D96W+ E239C+ Q249R+ P256T+ G266S E1SPPCGRRP (SEQ ID NO:25) + E239C+ Q249R+ P256T+ G266S+ T267A E1SPPCGRRP (SEQ ID NO:25) + E239C+ Q249R+ G266D E1SPPCGRRP (SEQ ID NO:25) + G91A+ D96W+ E239C+ Q249R+ G266D E1SPPRRP (SEQ ID NO:23) + D96S+ E239C+ Q249R+ G266D 12598 G266D G91A +D96W +F99K +G263Q +L264A +L265T +G266D +T267A +L269N +270A+ 271G+ 272G+ 273F (+274S) G266E G263A +G266A E1SPCRPRP (SEQ ID NO:26) +E239C +Q249R +G266A E1SPCRPRP (SEQ ID NO:26) +E239C +Q249R +G266S D96S + G266A D96S + G266S D96S + G266W E1SPPCGRRP (SEQ ID NO:23) +D96S +E239C +Q249R +G263D +L264I +I265N +G266E E1SPPCGRRP (SEQ ID NO:23) +D96S +E239C +Q249R +L264I +I265N +G266T +T267GL

D96F +G266A
D96F +G266S
E1SPPCGRRP (SEQ ID NO:23) +E99N +E239C +Q249R +G266A
E1SPPCGRRP (SEQ ID NO:23) + D96S +E239C +Q249R +G266A
E1SPPCGRRP (SEQ ID NO:23) + D96S +E239C +Q249R +G266S
E1SPPCGRRP (SEQ ID NO:23) + D96S +E239C +Q249R +G263F +L264A +G266S +T267E
V60G +D62A +S83T +R84K +D96W +G266D
V60G +D62A +S83T +D96W +G266D
V60G +D62A +S83T +D96W +G266W
L259I
L259N
D96W +G263Q +L264A +i265T +G266D +T267A

Please amend the tables beginning on page 45, below Example 7, as follows:

E1SPIRPRP (SEQ ID NO:22) +G91A +D96N +E99K +Q249R
E1SPCRPRP (SEQ ID NO:26) + S83T+ N94K+ D96L+ E239C+ Q249R
G266D
E1SPIRPRP (SEQ ID NO:22) +D62A +E99K +Q249R
E1SPIRPRP (SEQ ID NO:22) +D62G +E99K +Q249R
E1SPIRPRP (SEQ ID NO:22) +D62V +E99K +Q249R
E1SPIRPRP (SEQ ID NO:22) +R84W +E99K +Q249R
E1SPIRPRP (SEQ ID NO:22) +R84K +E99K +Q249R
E1SPIRPRP (SEQ ID NO:22) + K98D +E99K +Q249R
E1SPIRPRP (<u>SEQ ID NO:22)</u> + E99K +Q249R + 270PGLPFKRV
E1SPPCGRRP (SEQ ID NO:25) + E99N +N101S +T231K +R232G +D234G +E239C +Q249R
E1SPIRPRP (SEQ ID NO:22) +E99K +Q249R + 270PWPARLGRL
L93K +D96G
G91A +D96W +E99K +G263Q +L264A +I265T +G266D +T267A +L269N +270A+ 271G+ 272G+ 273F (+274S)
E1SPCRPRP +V60G +E99N +S119G +R209P +E239C +Q249R
G266A
G266E

G266V
G263Q +L264A +l265T +G266D +T267A
G266L
G263A +G266A
E1SPCRPRP (SEQ ID NO:26) +E239C +Q249R +G266A
E1SPCRPRP (SEQ ID NO:26) +E239C +Q249R +G266S
D96S + G266A
D96S + G266S
D96S + G266W
L264I +I265N +G266T +T267GL
E1SPPCGRRP (SEQ ID NO:25) +D96S +E239C +Q249R +L264I +I265N +G266T +T267GL
D96F +G266A
D96F +G266S
E1SPPCGRRP (SEQ ID NO:25) +E99N +E239C +Q249R +G266A
E1SPPCGRRP (SEQ ID NO:25) + D96S +E239C +Q249R +G266A
E1SPPCGRRP (SEQ ID NO:25) + D96S +E239C +Q249R +G266S
D62A + S83T
E1SPPCGRRP (SEQ ID NO:25) +K98D +E99N +E239C +Q249R
T231R +N233R +270CP
E1SPPCGRRP (SEQ ID NO:25) +E99N +E239C +Q249R +270MD
E1SPPCGRRP (SEQ ID NO:25) + D62A +S83T +E99N +E239C +Q249R
D62A +S83T + G91A +E99K +T231R +N233R +Q249R
V60G +D62A +S83T +R84K +D96W +G266D
L259N
L259R
L259M
L259Q
SPPCGRRP(-E) (<u>SEQ ID NO:25)</u> +R84W +E99N +N101S +E239C +Q249R
R84W +G91A +E99K +T231R +N233R +Q249R
Y21I
Y21V
SPIRPRP(-E) (SEQ ID NO:22) +R84L +E99K +Q249R
·

Y21C

SPIRPRP(-E) (SEQ ID NO:22) +D62 +E99K +Q249R

D96W+G263Q+L264A+l265T+G266D+T267A+L269N+A270+G271+G272+F273+S274.

G91A +D96W +E99K +G263Q +L264A +l265T +G266D +T267A +L269N +270A +271G +272G +273F +274S

The following variants of the parent lipase from *Humicola lanuginosa* may also have an increased specificity for long-chain fatty acids:

SPIRPRP(-E) (SEQ ID NO:22) +V60R +D62V +L93K +E99K +Q249R

SPIRPRP(-E) (SEQ ID NO:22) +D62V +E99K +Q249R

SPIRPRP(-E) (SEQ ID NO:22) +E99K +Q249R +P256D

SPIRPRP(-E) (SEQ ID NO:22) +D62V +E99K +Q249R +P256D

SPIRPRP(-E) (SEQ ID NO:22) +D62V +E99K +Q249R +P256S

G91A +D96W +E99K +G263Q +L264A +I265T +G266D +T267A +L269N +270A +271G +272G +273F +274S

G91A +D96L +E99K +G263Q +L264A +I265T +G266D +T267A +L269N +270A +271G +272G +273F +274S

G91A +D96N +E99K +G263Q +L264A +I265T +G266D +T267A +L269N +270A +271G +272G +273F +274S

G91A +D96A +E99K +G263Q +L264A +I265T +G266D +T267A +L269N +270A +271G +272G +273F +274S

G91A +D96E +E99K +G263Q +L264A +I265T +G266D +T267A +L269N +270A +271G +272G +273F +274S

G91A +D96S +E99K +G263Q +L264A +I265T +G266D +T267A +L269N +270A +271G +272G +273F +274S

G91A +D96R +E99K +G263Q +L264A +I265T +G266D +T267A +L269N +270A +271G +272G +273F +274S

G91A +D96G +E99K +G263Q +L264A +I265T +G266D +T267A +L269N +270A +271G +272G +273F +274S

G91A +D96Q +E99K +G263Q +L264A +I265T +G266D +T267A +L269N +270A +271G +272G +273F +274S

G91A +D96F +E99K +G263Q +L264A +I265T +G266D +T267A +L269N +270A +271G +272G +273F +274S

G91A +D96W +E99K +G263Q +L264A +I265T +G266S +T267A +L269N +270A +271G +272G +273F +274S

G91A +D96F +E99K +G263Q +L264A +I265T +G266S +T267A +L269N +270A +271G +272G +273F +274S

R84W +G91A +D96W +E99K +G263Q +L264A +I265T +G266S +T267A +L269N +270A

+271G +272G +273F +274S

R84W +G91A +D96F +E99K +G263Q +L264A +l265T +G266S +T267A +L269N +270A +271G +272G +273F +274S

R84W+G91A+D96F+E99K+G263Q+L264A+I265T+G266D+T267A+L269N+270A+271G +272G+273F+274S

SPPCGRRP(-E) (SEQ_ID_NO:25) +V60G +D62E +S83T +R84K +E99N +N101S +E239C +Q249R

V60G +D62E +S83T +R84K +G91A +E99K +T231R +N233R +Q249R

SPIRPRP(-E) (SEO ID NO:22) +E99K +R195O +R209E +O249R

Please amend the table on page 49, below Example 10, as follows:

3FIRFRF(-E) [3EQ ID 140.22] +E99K +R 193Q +R209E +Q249K
N101R +R195Q +R209E +L259S +Y261D
N101R +R195Q +R209E +L259S
N101R +L259S +Y261D
N101R +L259S
Y261D
L259S
SPIRPRP(-E) (SEQ ID NO:22) +E99K +N101R +Q249R
G263D +L264I +I265N +G266E +T267GS
Y261I
D234R
Y261K